

STN Columbus

* * * * * * * * * * Welcome to STN International * * * * * * * * * * *

| | | | |
|--------------|----|--------|--|
| NEWS | 1 | | Web Page URLs for STN Seminar Schedule - N. America |
| NEWS | 2 | | "Ask CAS" for self-help around the clock |
| NEWS | 3 | FEB 25 | CA/CAPLUS - Russian Agency for Patents and Trademarks (ROSPATENT) added to list of core patent offices covered |
| NEWS | 4 | FEB 28 | PATDPAFULL - New display fields provide for legal status data from INPADOC |
| NEWS | 5 | FEB 28 | BABS - Current-awareness alerts (SDIs) available |
| NEWS | 6 | FEB 28 | MEDLINE/LMEDLINE reloaded |
| NEWS | 7 | MAR 02 | GBFULL: New full-text patent database on STN |
| NEWS | 8 | MAR 03 | REGISTRY/ZREGISTRY - Sequence annotations enhanced |
| NEWS | 9 | MAR 03 | MEDLINE file segment of TOXCENTER reloaded |
| NEWS | 10 | MAR 22 | KOREPAT now updated monthly; patent information enhanced |
| NEWS | 11 | MAR 22 | Original IDE display format returns to REGISTRY/ZREGISTRY |
| NEWS | 12 | MAR 22 | PATDPASPC - New patent database available |
| NEWS | 13 | MAR 22 | REGISTRY/ZREGISTRY enhanced with experimental property tags |
| NEWS | 14 | APR 04 | EPFULL enhanced with additional patent information and new fields |
| NEWS | 15 | APR 04 | EMBASE - Database reloaded and enhanced |
| NEWS | 16 | APR 18 | New CAS Information Use Policies available online |
| NEWS | 17 | APR 25 | Patent searching, including current-awareness alerts (SDIs), based on application date in CA/CAplus and USPATFULL/USPAT2 may be affected by a change in filing date for U.S. applications. |
| NEWS | 18 | APR 28 | Improved searching of U.S. Patent Classifications for U.S. patent records in CA/CAplus |
| NEWS EXPRESS | | | JANUARY 10 CURRENT WINDOWS VERSION IS V7.01a, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), AND CURRENT DISCOVER FILE IS DATED 10 JANUARY 2005 |
| NEWS HOURS | | | STN Operating Hours Plus Help Desk Availability |
| NEWS INTER | | | General Internet Information |
| NEWS LOGIN | | | Welcome Banner and News Items |
| NEWS PHONE | | | Direct Dial and Telecommunication Network Access to STN |
| NEWS WWW | | | CAS World Wide Web Site (general information) |

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FILE 'HOME' ENTERED AT 13:28:13 ON 18 MAY 2005

=> fil reg
COST IN U.S. DOLLARS

FULL ESTIMATED COST

| SINCE FILE ENTRY | TOTAL SESSION |
|---------------------|------------------|
| 0.21 | 0.21 |

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FILE 'REGISTRY' ENTERED AT 13:28:19 ON 18 MAY 2005
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Property values tagged with IC are from the ZIC/VINITI data file
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STRUCTURE FILE UPDATES: 17 MAY 2005 HIGHEST RN 850605-77-5
DICTIONARY FILE UPDATES: 17 MAY 2005 HIGHEST RN 850605-77-5

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

Please note that search-term pricing does apply when
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*
* The CA roles and document type information have been removed from *
* the IDE default display format and the ED field has been added, *
* effective March 20, 2005. A new display format, IDERL, is now *
* available and contains the CA role and document type information. *
*

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more
information enter HELP PROP at an arrow prompt in the file or refer
to the file summary sheet on the web at:
<http://www.cas.org/ONLINE/DBSS/registryss.html>

=> s eplerenone/cn
L1 1 EPLERENONE/CN

=> d

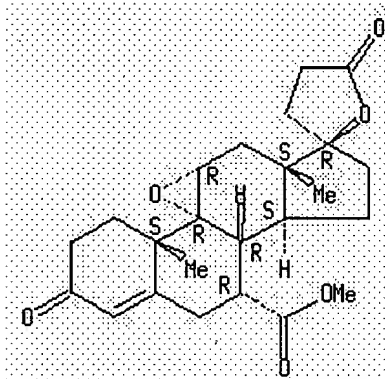
L1 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2005 ACS on STN
RN 107724-20-9 REGISTRY
ED Entered STN: 26 Apr 1987
CN Pregn-4-ene-7,21-dicarboxylic acid, 9,11-epoxy-17-hydroxy-3-oxo-,
γ-lactone, methyl ester, (7α,11α,17α)- (9CI) (CA
INDEX NAME)
OTHER CA INDEX NAMES:
CN Spiro[9,11-epoxy-9H-cyclopenta[a]phenanthrene-17(2H),2'(3'H)-furan],
pregn-4-ene-7,21-dicarboxylic acid deriv.

OTHER NAMES:
CN CGP 30083
CN Eplerenone
CN Epoxymexrenone
CN Inspira
CN SC 6110
CN SC 66110
FS STEREOSEARCH
MF C24 H30 O6
CI COM
SR CA
LC STN Files: ADISINSIGHT, ADISNEWS, ANABSTR, BIOSIS, BIOTECHNO, CA,

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CAPLUS, CASREACT, CIN, DDFU, DIOGENES, DRUGU, EMBASE, IMSDRUGNEWS,
IMSPATENTS, IMSRESEARCH, IPA, MEDLINE, MRCK*, PATDPASPC, PHAR, PROMT,
PROUSDDR, PS, SYNTHLINE, TOXCENTER, USAN, USPAT2, USPATFULL
(*File contains numerically searchable property data)

Absolute stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

190 REFERENCES IN FILE CA (1907 TO DATE)
4 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
192 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> s eplerenone
L2 1 EPLERENONE

=> fil medl Hcapl biosis Uspatf wpids
COST IN U.S. DOLLARS SINCE FILE TOTAL
SESSION
FULL ESTIMATED COST 11.47 11.68

FILE 'MEDLINE' ENTERED AT 13:29:07 ON 18 MAY 2005

FILE 'HCAPLUS' ENTERED AT 13:29:07 ON 18 MAY 2005
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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FILE 'BIOSIS' ENTERED AT 13:29:07 ON 18 MAY 2005
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FILE 'USPATFULL' ENTERED AT 13:29:07 ON 18 MAY 2005
CA INDEXING COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'WPIDS' ENTERED AT 13:29:07 ON 18 MAY 2005
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=> s 12

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COMMAND INTERRUPTED

4 FILES SEARCHED...

If this message appears repeatedly, please notify the Help Desk.
 Enter "HELP STN" for information on contacting the nearest STN Help
 Desk by telephone or via SEND in the STNMAIL file.

=> s 12

L3 499 L2

=> s heart failure

L4 168243 HEART FAILURE

=> s 13 and 14

L5 235 L3 AND L4

=> s 13 (S) 14

L6 30 L3 (S) L4

=> d ibib abs 27-30

L6 ANSWER 27 OF 30 WPIDS COPYRIGHT 2005 THE THOMSON CORP on STN
Full Text

ACCESSION NUMBER: 2002-062616 [08] WPIDS
 CROSS REFERENCE: 2001-343553 [36]; 2002-055827 [07]; 2002-195909 [25];
 2002-227077 [28]; 2003-843079 [78]; 2003-864187 [80];
 2004-021401 [02]
 DOC. NO. CPI: C2002-017915
 TITLE: Treatment of pathogenic effects in patients with
 sub-normal endogenous aldosterone level, salt sensitivity
 and/or elevated sodium dietary intake comprises
 administration of epoxy-steroidal aldosterone
 antagonists.
 DERWENT CLASS: B01
 INVENTOR(S): FEDDE, K N; FUNDER, J W; GARTHWAITE, S M; ROCHA, R;
 RONIKER, B; WILLIAMS, G H
 PATENT ASSIGNEE(S): (PHAA) PHARMACIA CORP
 COUNTRY COUNT: 93
 PATENT INFORMATION:

| PATENT NO | KIND | DATE | WEEK | LA | PG |
|---------------|---|--------------------|--------|----|----|
| WO 2001095893 | A1 | 20011220 (200208)* | EN 317 | | |
| RW: | AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW | | | | |
| W: | AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW | | | | |
| AU 2001016580 | A | 20011224 (200227) | | | |

APPLICATION DETAILS:

| PATENT NO | KIND | APPLICATION | DATE |
|---------------|------|-----------------|----------|
| WO 2001095893 | A1 | WO 2000-US31263 | 20001114 |

FILING DETAILS:

| PATENT NO | KIND | PATENT NO |
|---------------|------------|---------------|
| AU 2001016580 | A Based on | WO 2001095893 |

PRIORITY APPLN. INFO: US 2000-233056P 20000914; US
 2000-211064P 20000613; US
 2000-211250P 20000613; US
 2000-211253P 20000613; US
 2000-211264P 20000613; US
 2000-211311P 20000613; US
 2000-211340P 20000613; US
 2000-211451P 20000613; US
 2000-211459P 20000613; US
 2000-221358P 20000727; US
 2000-221364P 20000727

AN 2002-062616 [08] WPIDS

CR 2001-343553 [36]; 2002-055827 [07]; 2002-195909 [25]; 2002-227077 [28];
2003-843079 [78]; 2003-864187 [80]; 2004-021401 [02]

AB WO 200195893 A UPAB: 20040107

NOVELTY - Treatment or prophylaxis of one or more pathogenic effects in a human comprises administration of one or more epoxy-steroidal aldosterone antagonists, where the patient has a sub-normal endogenous aldosterone level, salt sensitivity and/or elevated sodium dietary intake.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) a method for treatment or prophylaxis of heart failure in a human suffering from or susceptible to cardiovascular disease comprising administration of an angiotensin converting enzyme (ACE) inhibitor, a loop diuretic and one or more aldosterone antagonists, where the patient has salt sensitivity or elevated dietary sodium intake;

(2) a method for reducing or reversing the progression of salt sensitivity by administering one or more epoxy-steroidal aldosterone antagonists;

(3) a method of reducing or preventing one or more pathogenic effects resulting from aberrant aldosterone levels in the brain by administering one or more epoxy-steroidal aldosterone antagonists;

(4) a method for reducing or preventing one or more pathogenic effects resulting from aberrant sodium retention in the kidney by administering one or more epoxy-steroidal aldosterone antagonists;

(5) a method for treating salt-sensitive hypertension by administering eplerenone; and

(6) a method for treating salt-sensitive **heart failure** by administering **eplerenone**.

ACTIVITY - Cardiant; Vasotropic; Hypotensive; Nephrotropic;
Antidiabetic; Ophthalmological; Cerebroprotective; Hepatotropic;
Neuroprotective; Antimigraine; Antiinflammatory.

A test to demonstrate that eplerenone can prevent aldosterone/salt-mediated early cardiovascular injury to the heart was carried out in male Wistar rats. The model combined elevated blood pressure, moderately high blood salt intake, an activated renin-angiotensin-aldosterone system (RAAS) and suppressed nitric oxide production. The model involved inhibiting nitric oxide synthase with Nomega-nitro-L-arginine methyl ester (L-NAME) for 14 days in 1% NaCl-drinking rats combined with a 3-day infusion of angiotensin II on days 11-14. At the end of the experiment, cardiac hypertrophy index was higher in rats receiving L-NAME/angiotensin II/NaCl. Infusion of

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aldosterone reversed the effect of adrenalectomy on cardiac hypertrophy index.

MECHANISM OF ACTION - Aldosterone antagonist.

USE - For treating hypertension, cardiovascular disease, heart failure, vascular disease, renal dysfunction, liver disease, cerebrovascular disease, retinopathy, neuropathy, insulinopathy, edema, endothelial dysfunction, baroreceptor dysfunction, migraine, hot flashes, premenstrual tension and salt sensitivity (all claimed).

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L6 ANSWER 28 OF 30 WPIDS COPYRIGHT 2005 THE THOMSON CORP on STN

Full Text

ACCESSION NUMBER: 2001-451592 [48] WPIDS
 DOC. NO. CPI: C2001-136351
 TITLE: Nanoparticulate eplerenone, which is useful in treatment of hyperaldosteronism and related conditions, exhibits improved bioavailability and can be used at reduced dosages.
 DERWENT CLASS: A96 B01
 INVENTOR(S): DESAI, S; GOKHALE, R D; THOSAR, S S; TOLBERT, D S
 PATENT ASSIGNEE(S): (PHAA) PHARMACIA CORP; (DESA-I) DESAI S; (GOKH-I) GOKHALE R D; (THOS-I) THOSAR S S; (TOLB-I) TOLBERT D S
 COUNTRY COUNT: 95
 PATENT INFORMATION:

| PATENT NO | KIND | DATE | WEEK | LA | PG |
|---------------|--|--------------------|------|----|----|
| <hr/> | | | | | |
| WO 2001041770 | A2 | 20010614 (200148)* | EN | 64 | |
| RW: | AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW | | | | |
| W: | AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW | | | | |
| AU 2001017562 | A | 20010618 (200161) | | | |
| US 2002006919 | A1 | 20020117 (200212) | | | |
| EP 1175220 | A2 | 20020130 (200216) | EN | | |
| R: | AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR | | | | |
| US 2003212053 | A1 | 20031113 (200382) | | | |
| EP 1527782 | A1 | 20050504 (200530) | EN | | |
| R: | AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE TR | | | | |

APPLICATION DETAILS:

| PATENT NO | KIND | APPLICATION | DATE |
|---------------|----------------|-----------------|----------|
| <hr/> | | | |
| WO 2001041770 | A2 | WO 2000-US30179 | 20001204 |
| AU 2001017562 | A | AU 2001-17562 | 20001204 |
| US 2002006919 | A1 Provisional | US 1999-169658P | 19991208 |
| | Provisional | US 2000-208981P | 20000602 |
| | | US 2000-732246 | 20001207 |
| EP 1175220 | A2 | EP 2000-980277 | 20001204 |
| | | WO 2000-US30179 | 20001204 |
| US 2003212053 | A1 Provisional | US 1999-169658P | 19991208 |
| | Provisional | US 2000-208981P | 20000602 |
| | Div ex | US 2000-732246 | 20001207 |
| | | US 2003-417602 | 20030416 |
| EP 1527782 | A1 Div ex | EP 2000-980277 | 20001204 |
| | | EP 2004-30120 | 20001204 |

FILING DETAILS:

| PATENT NO | KIND | PATENT NO |
|---|---|--|
| AU 2001017562 | A Based on | WO 2001041770 |
| EP 1175220 | A2 Based on | WO 2001041770 |
| EP 1527782 | A1 Div ex | EP 1175220 |
| PRIORITY APPLN. INFO: US 2000-208981P 1999-169658P 2000-732246 2003-417602 | | 20000602; US 19991208; US 20001207; US 20030416 |
| AN | 2001-451592 [48] | WPIDS |
| AB | WO 200141770 A | UPAB: 20010829 |
| NOVELTY - Eplerenone particles, in which 90 wt.% of the particles are smaller than 15 microns, are new. | | |
| DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for a pharmaceutical composition comprising 10-1,000 mg of the eplerenone particles and one or more excipients. | | |
| ACTIVITY - Hypotensive; Cardiant; Cerebroprotective; Vasotropic; Hepatotropic. | | |
| MECHANISM OF ACTION - Aldosterone receptor antagonist. | | |
| USE - Eplerenone (9,11 alpha -epoxy-17 alpha -3-oxopregn-4-ene-7 alpha ,21-dicarboxylate, gamma -lactone) is an aldosterone receptor antagonist which can be used in treatment of conditions associated with hyperaldosteronism, e.g., heart failure , cardiac insufficiency, hypertension, edema associated with liver insufficiency, post-myocardial infarction, cirrhosis of the liver and accelerated heart rate. It can be used to prevent stroke. | | |
| ADVANTAGE - Reducing the particle size of solid eplerenone can improve the bioavailability of eplerenone and reduce the amount of eplerenone administered. Eplerenone interacts minimally with steroid receptors other than aldosterone receptors, e.g. progestin, androgen or glucocorticoid receptors. | | |
| Dwg. 0/0 | | |
| L6 | ANSWER 29 OF 30 | WPIDS COPYRIGHT 2005 THE THOMSON CORP on STN |
| <u>Full Text</u> | | |
| ACCESSION NUMBER: | 2001-441528 [47] WPIDS | |
| CROSS REFERENCE: | 1997-341331 [31]; 1998-413631 [35]; 2001-464975 [50] | |
| DOC. NO. CPI: | C2001-133333 | |
| TITLE: | New Form L crystalline eplerenone, useful for treating or preventing aldosterone-mediated conditions such as hypertension. | |
| DERWENT CLASS: | B01 D16 | |
| INVENTOR(S): | BARTON, K P; BORSCHARDT, T B; CARLOS, M V; DESAI, S; FERRO, L J; GANSER, S; GAUD, H T; LITTLE, C R; MUDIPALLI, P S; PIETZ, M A; PILIPAUASKAS, D R; SING, Y L; STAHL, G L; WEICZOREK, J J; YAN, C Y; WIECZOREK, J J; BORCHARDT, T; BORCHARDT, T B; GANSER, S S | |
| PATENT ASSIGNEE(S): | (PHAA) PHARMACIA CORP; (BART-I) BARTON K P; (BORC-I) BORCHARDT T B; (CARL-I) CARLOS M V; (DESA-I) DESAI S; (FERR-I) FERRO L J; (GANS-I) GANSER S S; (GAUD-I) GAUD H T; (LITT-I) LITTLE C R; (MUDI-I) MUDIPALLI P S; (PIET-I) PIETZ M A; (PILI-I) PILIPAUASKAS D R; (SING-I) SING Y L; (STAHL-I) STAHL G L; (WIEC-I) WIECZOREK J J; (YANC-I) YAN C Y | |
| COUNTRY COUNT: | 87 | |
| PATENT INFORMATION: | | |

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| PATENT NO | KIND | DATE | WEEK | LA | PG |
|---------------|--|----------------------|------|-----|----|
| WO 2001041535 | A2 | 20010614 (200147)* | EN | 165 | |
| RW: | AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW | | | | |
| W: | AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZW | | | | |
| AU 2001020411 | A | 20010618 (200161) | | | |
| NO 2001003856 | A | 20011008 (200171) | | | |
| NO 2001003857 | A | 20011008 (200171) | | | |
| EP 1175434 | A2 | 20020130 (200216) EN | | | |
| R: | AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR | | | | |
| BR 2000008054 | A | 20020312 (200226) | | | |
| US 2002045746 | A1 | 20020418 (200228) | | | |
| KR 2001112261 | A | 20011220 (200239) | | | |
| KR 2002003192 | A | 20020110 (200247) | | | |
| HU 2002001457 | A2 | 20020828 (200264) | | | |
| ZA 2001007147 | A | 20030226 (200321) | 176 | | |
| JP 2003515611 | W | 20030507 (200331) | 183 | | |
| US 2003083493 | A1 | 20030501 (200331) | | | |
| CN 1433427 | A | 20030730 (200365) | | | |
| NZ 513962 | A | 20040827 (200460) | | | |
| AU 2004242560 | A1 | 20050127 (200525) # | | | |

APPLICATION DETAILS:

| PATENT NO | KIND | APPLICATION | DATE |
|---------------|----------------|-----------------|----------|
| WO 2001041535 | A2 | WO 2000-US30178 | 20001204 |
| AU 2001020411 | A | AU 2001-20411 | 20001204 |
| NO 2001003856 | A | WO 2000-US32416 | 20001204 |
| | | NO 2001-3856 | 20010808 |
| NO 2001003857 | A | WO 2000-US30178 | 20001204 |
| | | NO 2001-3857 | 20010808 |
| EP 1175434 | A2 | EP 2000-983683 | 20001204 |
| | | WO 2000-US30178 | 20001204 |
| BR 2000008054 | A | BR 2000-8054 | 20001204 |
| | | WO 2000-US30178 | 20001204 |
| US 2002045746 | A1 Provisional | US 1995-8455P | 19951211 |
| | Div ex | US 1996-763910 | 19961211 |
| | Provisional | US 1997-49388P | 19970611 |
| | CIP of | WO 1997-US23090 | 19971211 |
| | CIP of | US 1999-246204 | 19990208 |
| | CIP of | US 1999-246908 | 19990209 |
| | Provisional | US 1999-169556P | 19991208 |
| | Provisional | US 1999-169608P | 19991208 |
| | Provisional | US 1999-169639P | 19991208 |
| | Provisional | US 1999-169683P | 19991208 |
| | Provisional | US 1999-169707P | 19991208 |
| | Provisional | US 1999-169807P | 19991208 |
| | CIP of | US 1999-319673 | 19991213 |
| | CIP of | US 2000-583137 | 20000530 |
| | CIP of | US 2000-583158 | 20000530 |
| | | US 2000-732209 | 20001207 |
| KR 2001112261 | A | KR 2001-710043 | 20010808 |
| KR 2002003192 | A | KR 2001-710042 | 20010808 |

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|---------------|----------------|-----------------|----------|
| HU 2002001457 | A2 | WO 2000-US30178 | 20001204 |
| | | HU 2002-1457 | 20001204 |
| ZA 2001007147 | A | ZA 2001-7147 | 20010829 |
| JP 2003515611 | W | WO 2000-US30178 | 20001204 |
| | | JP 2001-542722 | 20001204 |
| US 2003083493 | A1 Provisional | US 1999-169556P | 19991208 |
| | Provisional | US 1999-169608P | 19991208 |
| | Provisional | US 1999-169639P | 19991208 |
| | Provisional | US 1999-169683P | 19991208 |
| | Provisional | US 1999-169707P | 19991208 |
| | Provisional | US 1999-169807P | 19991208 |
| | Cont of | US 2000-732209 | 20001207 |
| | | US 2002-191626 | 20020709 |
| CN 1433427 | A | CN 2000-805771 | 20001204 |
| NZ 513962 | A | NZ 2000-513962 | 20001204 |
| | | WO 2000-US30178 | 20001204 |
| AU 2004242560 | A1 Div ex | AU 2001-20411 | 20001204 |
| | | AU 2004-242560 | 20041231 |

FILING DETAILS:

| PATENT NO | KIND | PATENT NO |
|---------------|-------------|---------------|
| AU 2001020411 | A Based on | WO 2001041535 |
| EP 1175434 | A2 Based on | WO 2001041535 |
| BR 2000008054 | A Based on | WO 2001041535 |
| US 2002045746 | A1 Div ex | US 5981744 |
| | Div ex | US 6180780 |
| | CIP of | US 6258946 |
| HU 2002001457 | A2 Based on | WO 2001041535 |
| JP 2003515611 | W Based on | WO 2001041535 |
| NZ 513962 | A Div in | NZ 533700 |
| | Based on | WO 2001041535 |

PRIORITY APPLN. INFO: US 1999-169807P 19991208; US
 1999-169556P 19991208; US
 1999-169608P 19991208; US
 1999-169639P 19991208; US
 1999-169683P 19991208; US
 1999-169707P 19991208; US
 1999-169682P 19991208; US
 1999-169690P 19991208; US
 1995-8455P 19951211; US
 1996-763910 19961211; US
 1997-49388P 19970611; WO
 1997-US23090 19971211; US
 1999-246204 19990208; US
 1999-246908 19990209; US
 1999-319673 19991213; US
 2000-583137 20000530; US
 2000-583158 20000530; US
 2000-732209 20001207; US
 2002-191626 20020709; AU
 2004-242560 20041231

AN 2001-441528 [47] WPIDS
 CR 1997-341331 [31]; 1998-413631 [35]; 2001-464975 [50]
 AB WO 200141535 A UPAB: 20050419

NOVELTY - A new Form L crystalline eplerenone is disclosed.

DETAILED DESCRIPTION - (A) Form L crystalline eplerenone having a monoclinic crystal system and an X-ray powder diffraction pattern with a

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peak at 8.0 plus or minus 0.2 deg. 2 Omega is claimed.

An INDEPENDENT CLAIM is also included for an eplerenone drug substance comprising Form L crystalline eplerenone.

ACTIVITY - Hypotensive; Cardiant; Hepatotropic; Cytostatic; Antidepressant.

MECHANISM OF ACTION - Aldosterone receptor antagonist.

USE - The Form L crystalline **eplerenone** can be used for treating or preventing an aldosterone-mediated condition or disorder (claimed). It can be used for the treatment of conditions associated with hyperaldosteronism such as hypertension, **heart failure** including cardiac insufficiency, cirrhosis of the liver, excess collagen, fibrosis, benign prostate hypertrophy or depression.

ADVANTAGE - The new crystalline form has a high degree of physical stability at normal temperatures of storage and use. It can be used with other forms of eplerenone to provide compositions having a variety of dissolution profiles, including controlled-release compositions..

Dwg. 0/26

L6 ANSWER 30 OF 30 WPIDS COPYRIGHT 2005 THE THOMSON CORP on STN

Full Text

ACCESSION NUMBER: 2000-524152 [47] WPIDS
 DOC. NO. CPI: C2000-155613
 TITLE: Novel composition containing 10 - 100 mg **eplerenone**, useful for the treatment of e.g. **heart failure**, hypertension and post-myocardial infarction, exhibits superior activity, potency and safety than prior art.
 DERWENT CLASS: A96 B02 C02
 INVENTOR(S): GOKHALE, R D; THOSAR, S S; TOLBERT, D S
 PATENT ASSIGNEE(S): (SEAR) SEARLE & CO G D; (GOKH-I) GOKHALE R D; (THOS-I) THOSAR S S; (TOLB-I) TOLBERT D S
 COUNTRY COUNT: 91
 PATENT INFORMATION:

| PATENT NO | KIND DATE | WEEK | LA | PG |
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| <hr/> | | | | |
| WO 2000033847 | A1 20000615 (200047)* | EN 196 | | |
| RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ TZ UG ZW | | | | |
| W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW | | | | |
| AU 2000019368 | A 20000626 (200047) | | | |
| NO 2001002782 | A 20010703 (200154) | | | |
| EP 1135139 | A1 20010926 (200157) EN | | | |
| R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI | | | | |
| BR 9915964 | A 20010828 (200158) | | | |
| CZ 2001001942 | A3 20011212 (200206) | | | |
| CN 1329494 | A 20020102 (200227) | | | |
| KR 2001101132 | A 20011114 (200230) | | | |
| MX 2001005647 | A1 20010801 (200238) | | | |
| US 6410054 | B1 20020625 (200246) | | | |
| HU 2001004718 | A2 20020528 (200249) | | | |
| US 2002136775 | A1 20020926 (200265) | | | |
| ZA 2001004361 | A 20020731 (200271) | 205 | | |
| JP 2002531508 | W 20020924 (200278) | 171 | | |
| US 6495165 | B1 20021217 (200307) | | | |
| US 6534093 | B1 20030318 (200322) | | | |
| US 2003072808 | A1 20030417 (200329) | | | |

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|--|----|----------|-------------|
| US 6558707 | B1 | 20030506 | (200338) |
| NZ 511869 | A | 20030530 | (200341) |
| US 6592902 | B2 | 20030715 | (200348) |
| AU 763166 | B | 20030717 | (200356) |
| EP 1135139 | B1 | 20030910 | (200360) EN |
| R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT RO SE | | | |
| DE 69911240 | E | 20031016 | (200376) |
| US 2003215518 | A1 | 20031120 | (200377) |
| ES 2207977 | T3 | 20040601 | (200437) |
| US 2004192661 | A1 | 20040930 | (200465) |
| US 6863902 | B2 | 20050308 | (200518) |

APPLICATION DETAILS:

| PATENT NO | KIND | APPLICATION | DATE | |
|---------------|------|-----------------|-----------------|----------|
| WO 2000033847 | A1 | WO 1999-US29136 | 19991208 | |
| AU 2000019368 | A | AU 2000-19368 | 19991208 | |
| NO 2001002782 | A | WO 1999-US29136 | 19991208 | |
| | | NO 2001-2782 | 20010606 | |
| EP 1135139 | A1 | EP 1999-963052 | 19991208 | |
| | | WO 1999-US29136 | 19991208 | |
| BR 9915964 | A | BR 1999-15964 | 19991208 | |
| | | WO 1999-US29136 | 19991208 | |
| CZ 2001001942 | A3 | WO 1999-US29136 | 19991208 | |
| | | CZ 2001-1942 | 19991208 | |
| CN 1329494 | A | CN 1999-814110 | 19991208 | |
| KR 2001101132 | A | KR 2001-707041 | 20010605 | |
| MX 2001005647 | A1 | MX 2001-5647 | 20010605 | |
| US 6410054 | B1 | Provisional | US 1998-111646P | 19981209 |
| | | | US 1999-456614 | 19991208 |
| HU 2001004718 | A2 | WO 1999-US29136 | 19991208 | |
| | | HU 2001-4718 | 19991208 | |
| US 2002136775 | A1 | Provisional | US 1998-111646P | 19981209 |
| | | Cont of | US 1999-456614 | 19991208 |
| | | | US 2002-66360 | 20020319 |
| ZA 2001004361 | A | ZA 2001-4361 | 20010528 | |
| JP 2002531508 | W | WO 1999-US29136 | 19991208 | |
| | | JP 2000-586339 | 19991208 | |
| US 6495165 | B1 | Provisional | US 1998-111646P | 19981209 |
| | | Cont of | US 1999-456614 | 19991208 |
| | | | US 2002-101361 | 20020319 |
| US 6534093 | B1 | Provisional | US 1998-111646P | 19981209 |
| | | Cont of | US 1999-456614 | 19991208 |
| | | | US 2002-100930 | 20020319 |
| US 2003072808 | A1 | Provisional | US 1998-111646P | 19981209 |
| | | Cont of | US 1999-456614 | 19991208 |
| | | | US 2002-100712 | 20020319 |
| US 6558707 | B1 | Provisional | US 1998-111646P | 19981209 |
| | | Cont of | US 1999-456614 | 19991208 |
| | | | US 2002-100712 | 20020319 |
| NZ 511869 | A | NZ 1999-511869 | 19991208 | |
| | | WO 1999-US29136 | 19991208 | |
| US 6592902 | B2 | Provisional | US 1998-111646P | 19981209 |
| | | Cont of | US 1999-456614 | 19991208 |
| | | | US 2002-66360 | 20020319 |
| AU 763166 | B | AU 2000-19368 | 19991208 | |
| EP 1135139 | B1 | EP 1999-963052 | 19991208 | |
| | | | WO 1999-US29136 | 19991208 |
| DE 69911240 | E | DE 1999-611240 | 19991208 | |

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|---------------|----------------|-----------------|----------|
| | | EP 1999-963052 | 19991208 |
| | | WO 1999-US29136 | 19991208 |
| US 2003215518 | A1 Provisional | US 1998-111646P | 19981209 |
| | Cont of | US 1999-456614 | 19991208 |
| | Cont of | US 2002-100930 | 20020319 |
| | | US 2002-289025 | 20021106 |
| ES 2207977 | T3 | EP 1999-963052 | 19991208 |
| US 2004192661 | A1 Provisional | US 1998-111646P | 19981209 |
| | Cont of | US 1999-456614 | 19991208 |
| | Cont of | US 2002-100930 | 20020319 |
| | Cont of | US 2002-289025 | 20021106 |
| | | US 2004-817577 | 20040402 |
| US 6863902 | B2 Provisional | US 1998-111646P | 19981209 |
| | Cont of | US 1999-456614 | 19991208 |
| | Div ex | US 2002-100930 | 20020319 |
| | | US 2002-289025 | 20021106 |

FILING DETAILS:

| PATENT NO | KIND | PATENT NO |
|---------------|------------------|---------------|
| AU 2000019368 | A Based on | WO 2000033847 |
| EP 1135139 | A1 Based on | WO 2000033847 |
| BR 9915964 | A Based on | WO 2000033847 |
| CZ 2001001942 | A3 Based on | WO 2000033847 |
| HU 2001004718 | A2 Based on | WO 2000033847 |
| JP 2002531508 | W Based on | WO 2000033847 |
| US 6495165 | B1 Cont of | US 6410054 |
| US 6534093 | B1 Cont of | US 6410054 |
| US 2003072808 | A1 Cont of | US 6410054 |
| US 6558707 | B1 Cont of | US 6410054 |
| NZ 511869 | A Based on | WO 2000033847 |
| AU 763166 | B Previous Publ. | AU 2000019368 |
| | Based on | WO 2000033847 |
| EP 1135139 | B1 Based on | WO 2000033847 |
| DE 69911240 | E Based on | EP 1135139 |
| | Based on | WO 2000033847 |
| US 2003215518 | A1 Cont of | US 6410054 |
| | Cont of | US 6534093 |
| ES 2207977 | T3 Based on | EP 1135139 |
| US 2004192661 | A1 Cont of | US 6410054 |
| | Cont of | US 6534093 |
| US 6863902 | B2 Cont of | US 6410054 |
| | Div ex | US 6534093 |

PRIORITY APPLN. INFO: US 1998-111646P 19981209; US
 1999-456614 19991208; US
 2002-66360 20020319; US
 2002-101361 20020319; US
 2002-100930 20020319; US
 2002-100712 20020319; US
 2002-289025 20021106; US
 2004-817577 20040402

AN 2000-524152 [47] WPIDS

AB WO 200033847 A UPAB: 20000925

NOVELTY - A composition (I) comprising 10 - 1000 mg eplerenone and a carrier is new.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is provided for the preparation of (I).

ACTIVITY - Hypotensive; cardioactive; antiinflammatory.

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The efficacy of eplerenone in the treatment of hypertension was evaluated in a multi-center, randomized, double-blind, placebo-lead-in, parallel group study. A treatment regimen of 200 mg eplerenone BID (not defined) resulted in a decrease in diastolic blood pressure of 9.4 mm Hg, compared to 1.00 for the placebo.

MECHANISM OF ACTION - Aldosterone receptor antagonist (claimed).

USE - (I) causes an average increase in blood serum renin concentrations over an interval of 12 - 24 hours after ingestion of at least 10 %, an increase in blood serum aldosterone concentrations of at least 50 % and an average decrease in diastolic blood pressure of at least 5 % (claimed). The composition is used to treat heart failure, hypertension, edema associated with liver insufficiency and post-myocardial infarction (claimed).

ADVANTAGE - (I) exhibits superior activity, potency, safety and therapeutic effectiveness than prior art.

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L3 ANSWER 499 OF 499 WPIDS COPYRIGHT 2005 THE THOMSON CORP on STN

Full Text

ACCESSION NUMBER: 2000-317820 [27] WPIDS
DOC. NO. CPI: C2000-096186
TITLE: Treating weight loss due to cachexia which occurs in liver cirrhosis, cardiac cachexia involves administering an agent which reduces sympathetic nervous system activity and/or improves cardiovascular reflex status.
DERWENT CLASS: B05
INVENTOR(S): ANKER, S D; COATS, A J S
PATENT ASSIGNEE(S): (IMCO-N) IMPERIAL COLLEGE INNOVATIONS LTD
COUNTRY COUNT: 21
PATENT INFORMATION:

| PATENT NO | KIND | DATE | WEEK | LA | PG |
|---|------|--------------------|------|----|----|
| ----- | | | | | |
| WO 2000021509 | A2 | 20000420 (200027)* | EN | 72 | |
| RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE | | | | | |
| W: JP US | | | | | |
| EP 1121111 | A2 | 20010808 (200146) | EN | | |
| R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE | | | | | |
| JP 2002527378 | W | 20020827 (200271) | | 74 | |

APPLICATION DETAILS:

| PATENT NO | KIND | APPLICATION | DATE |
|---------------|------|----------------|----------|
| ----- | | | |
| WO 2000021509 | A2 | WO 1999-GB3302 | 19991015 |
| EP 1121111 | A2 | EP 1999-947762 | 19991015 |
| | | WO 1999-GB3302 | 19991015 |
| JP 2002527378 | W | WO 1999-GB3302 | 19991015 |
| | | JP 2000-575485 | 19991015 |

FILING DETAILS:

| PATENT NO | KIND | PATENT NO |
|---------------|-------------|---------------|
| ----- | | |
| EP 1121111 | A2 Based on | WO 2000021509 |
| JP 2002527378 | W Based on | WO 2000021509 |

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PRIORITY APPLN. INFO: GB 1999-17181 19990723; GB
 1998-22458 19981015; GB
 1998-22459 19981015

AN 2000-317820 [27] WPIDS

AB WO 200021509 A UPAB: 20000606

NOVELTY - Treating weight loss due to an underlying disease in a patient comprises administering an agent (I) which reduces sympathetic nervous system activity and/or improves cardiovascular reflex status.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) treating weight loss due to an underlying disease in a patient which involves administering any one of the following compounds (III): a compound which inhibits the effect of aldosterone such as an aldosterone antagonist; a chymase inhibitor; a cathepsin inhibitor; a beta receptor blocker; an imidazoline receptor antagonist; a centrally acting alpha receptor antagonist; a peripherally acting alpha receptor antagonist; a ganglion blocking agent; a drug that has an effect on cardiovascular reflexes and thereby reduce SNS activity such as an opiate; scopolamine; endothelin receptor antagonist; a xanthine oxidase inhibitor; or erythropoietin;

(2) treating weight loss due to underlying disease in a patient which involves electrically stimulating the patient's muscles;

(3) use of the above mentioned compounds in the manufacture of a medicament for preventing or treating weight loss due to underlying disease (idiopathic cachexia) or aging in a patient and also for manufacture of an agent for enhancing exercise performance in a healthy individual;

(4) preventing or treating weight loss due to the aging process in the patient which involves administering an agent (I) which reduces sympathetic nervous system activity, or (III);

(5) preventing or treating weight loss due to aging process which involves electrically stimulating the patient's muscles;

(6) enhancing exercise performance in a patient by administering (I) which reduces sympathetic nervous system activity, or (III);

(7) enhancing exercise performance in a patient by which involves electrically stimulating the patient's muscles;

(8) preventing weight loss consequent to a cardiovascular disorder in a patient at risk of heart disease which involves administering (III) with an inhibiting effect on aldosterone; a beta receptor blocker; an imidazoline receptor antagonist; a centrally acting alpha receptor agonist; a peripherally acting alpha receptor antagonist; or a ganglion blocking agent.

ACTIVITY - Anabolic.

The biological activity of (III) was tested in human cachexia patients. A patient with cachexia due to chronic heart failure (CHF) and a second patient with CHF and a muscle myopathy suffering from idiopathic cachexia were treated with 50 mg of an angiotensin II receptor antagonist (losartan) daily. The clinical status and parameters of body composition, strength and treadmill exercise capacity at base line and during follow-up was study for 126 days in patient 1 and 83 days in patient 2. The results showed that in both patients the exercise capacity was improved during the study. Also patient 1 had a weight gain of 4.6 kg.

MECHANISM OF ACTION - Sympathetic nervous system activator; Cardiac reflex status enhancer (claimed).

USE - The method is useful for treating weight loss in a patient having idiopathic cachexia with an underlying disease such as AIDS, liver cirrhosis, chronic obstructive pulmonary disease with or without emphysema, chronic renal failure, chronic infections, cancer, heart disease including hypertension and chronic heart failure (claimed).

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| => fil reg | | SINCE FILE | TOTAL |
| COST IN U.S. DOLLARS | | ENTRY | SESSION |
| FULL ESTIMATED COST | | 47.28 | 58.96 |

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DICTIONARY FILE UPDATES: 17 MAY 2005 HIGHEST RN 850605-77-5

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<http://www.cas.org/ONLINE/DBSS/registryss.html>

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L7 1 QUINAPRIL/CN

=> fil medl hcapl biosis uspatf wpids

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|----------------------|------------|---------|
| COST IN U.S. DOLLARS | SINCE FILE | TOTAL |
| FULL ESTIMATED COST | ENTRY | SESSION |
| | 5.03 | 63.99 |

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FILE 'USPATFULL' ENTERED AT 13:39:30 ON 18 MAY 2005

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CA INDEXING COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'WPIDS' ENTERED AT 13:39:30 ON 18 MAY 2005
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=> s 17
L8 1609 L7

=> d his

(FILE 'HOME' ENTERED AT 13:28:13 ON 18 MAY 2005)

FILE 'REGISTRY' ENTERED AT 13:28:19 ON 18 MAY 2005
L1 1 S EPLERENONE/CN
L2 1 S EPLERENONE

FILE 'MEDLINE, HCAPLUS, BIOSIS, USPATFULL, WPIDS' ENTERED AT 13:29:07 ON
18 MAY 2005
L3 499 S L2
L4 168243 S HEART FAILURE
L5 235 S L3 AND L4
L6 30 S L3 (S) L4

FILE 'REGISTRY' ENTERED AT 13:38:59 ON 18 MAY 2005
L7 1 S QUINAPRIL/CN

FILE 'MEDLINE, HCAPLUS, BIOSIS, USPATFULL, WPIDS' ENTERED AT 13:39:30 ON
18 MAY 2005
L8 1609 S L7

=> s 17 and 14
L9 295 L7 AND L4

=> s 17 (S) 14
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FIELD CODE - 'AND' OPERATOR ASSUMED 'L27 (S) L13'
L10 36 L7 (S) L4

=> d ibib abs 35-36

L10 ANSWER 35 OF 36 HCAPLUS COPYRIGHT 2005 ACS on STN
Full Text

ACCESSION NUMBER: 1991:400473 HCAPLUS
DOCUMENT NUMBER: 115:473
TITLE: Effects of quinapril, a new angiotensin converting enzyme inhibitor, on left ventricular failure and survival in the cardiomyopathic hamster. Hemodynamic, morphological, and biochemical correlates
AUTHOR(S): Haleen, Stephen J.; Weishaar, Ronald E.; Overhiser, Ronald W.; Bousley, Richard F.; Keiser, Joan A.; Rapundalo, Stephen R.; Taylor, David G.
CORPORATE SOURCE: Parke-Davis Pharm. Res. Div., Warner-Lambert Co., Ann Arbor, MI, 48105, USA
SOURCE: Circulation Research (1991), 68(5), 1302-12
CODEN: CIRUAL; ISSN: 0009-7330
DOCUMENT TYPE: Journal
LANGUAGE: English
AB The effect of chronic therapy with quinapril on the temporal progression of left ventricular failure and survival was assessed in the CHF 146 cardiomyopathic (CM) hamster, which is an idiopathic model of congestive

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heart failure. Age-matched Golden Syrian (GS) hamsters served as normal controls. Quinapril was administered in the drinking water at av. daily doses of 10.2, 112.4, and 222.4 mg/kg/day. In untreated CM hamsters, in vitro left ventricular performance progressively deteriorated with increasing age beginning at roughly 180 days. This decline in left ventricular performance was accompanied by a decrease in coronary flow and an increase in left ventricular vol. Administration of quinapril from 180 to 300 days of age prevented the decline of in vitro left ventricular contractile performance and coronary flow and also reduced the age-dependent increases in left ventricular vol. The cardioprotective effects of quinapril were obsd. at doses of 112.4 and 222.4 mg/kg/day but not at 10.2 mg/kg/day. Lung angiotensin-converting enzyme activity was significantly inhibited by quinapril in GS and CM hamsters at 240 and 300 days of age at all dose levels. In contrast, significant inhibition of ventricular angiotensin converting enzyme activity was obsd. consistently at doses of 112.4 and 222.4 mg/kg/day quinapril but not at 10.2 mg/kg/day. In the survival protocol, CM and GS hamsters were treated with vehicle or quinapril (100 mg/kg/day) from 180 to 522 days of age. During the initial 210 days of treatment (from 180 to 390 days of age) 78.3% of the vehicle-treated CM hamsters died compared with 27.7% of quinapril-treated CM hamsters. Quinapril increased the median survival of CM hamsters by 32.9% (112 days). Thus, chronic quinapril therapy exerts a significant cardioprotective effect and also increases survival.

L10 ANSWER 36 OF 36 HCAPLUS COPYRIGHT 2005 ACS on STN

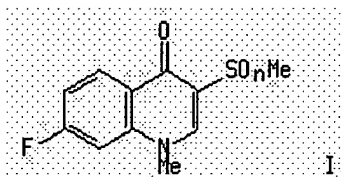
Full Text

ACCESSION NUMBER: 1991:199681 HCAPLUS
 DOCUMENT NUMBER: 114:199681
 TITLE: Treatment of heart failure with a quinolone derivative combined with an angiotensin-converting enzyme inhibitor
 INVENTOR(S): O'Connor, Patrick Coleman; Defesche, Charles Leon Marie
 PATENT ASSIGNEE(S): Boots Co. PLC, UK
 SOURCE: PCT Int. Appl., 27 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------|----------|-----------------|-------------|
| WO 9010445 | A1 | 19900920 | WO 1990-EP430 | 19900312 |
| W: AU, CA, JP, KR, US | | | | |
| RW: AT, BE, CH, DE, DK, ES, FR, GB, IT, LU, NL, SE | | | | |
| AU 9052616 | A1 | 19901009 | AU 1990-52616 | 19900312 |
| AU 640128 | B2 | 19930819 | | |
| JP 04503806 | T2 | 19920709 | JP 1990-504547 | 19900312 |
| EP 527720 | A1 | 19930224 | EP 1990-904349 | 19900312 |
| EP 527720 | B1 | 19940817 | | |
| R: AT, BE, CH, DE, DK, ES, FR, GB, IT, LI, LU, NL, SE | | | | |
| ES 2057542 | T3 | 19941016 | ES 1990-904349 | 19900312 |
| ZA 9001945 | A | 19910130 | ZA 1990-1945 | 19900314 |
| IL 93744 | A1 | 19950629 | IL 1990-93744 | 19900314 |
| PRIORITY APPLN. INFO.: | | | US 1989-324213 | A2 19890315 |
| | | | WO 1990-EP430 | A 19900312 |

OTHER SOURCE(S): MARPAT 114:199681

GI



AB The quinolone derivs. I ($n = 1, 2$) coadministered with angiotensin-converting enzyme inhibitors are more effective drugs for the treatment of heart failure, than the latter alone. Patients with congestive heart failure, that have received a daily dose 76.3 mg Captopril/day, and showed improvement of the hemodynamic parameters when the Captopril dose was reduced to 1/3 and Flosequinan (150 mg) was coadministered. Formulation examples are given.

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COST IN U.S. DOLLARS

| SINCE FILE | TOTAL |
|------------|---------|
| ENTRY | SESSION |

FULL ESTIMATED COST

14.56 78.55

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

| SINCE FILE | TOTAL |
|------------|---------|
| ENTRY | SESSION |

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